(FILE 'HOME' ENTERED AT 15:34:55 ON 21 JUL 2003)

INDEX 'ADISCTI, ADISINSIGHT, ADISNEWS, AGRICOLA, ANABSTR, AQUASCI, BIOBUSINESS, BIOCOMMERCE, BIOSIS, BIOTECHABS, BIOTECHDS, BIOTECHNO, CABA, CANCERLIT, CAPLUS, CEABA-VTB, CEN, CIN, CONFSCI, CROPB, CROPU, DDFB, DDFU, DGENE, DRUGB, DRUGLAUNCH, DRUGMONOG2, ...' ENTERED AT 15:35:36 ON 21 JUL 2003

SEA (ADIPONECTIN) OR (ADIPONECTIN-LIKE)

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      FILE SCISEARCH
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100
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      FILE USPATFULL
  6
      FILE WPIDS
     FILE WPINDEX
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FILE 'CAPLUS, SCISEARCH, MEDLINE, BIOSIS, EMBASE, ESBIOBASE, TOXCENTER, PASCAL, BIOTECHNO, JICST-EPLUS' ENTERED AT 15:37:00 ON 21 JUL 2003

L2 226 S L1 AND (ISOLAT? OR CHARACT? OR PURIF?)

19 S L2 AND PURIF?

L1

L3

L4

11 DUP REM L3 (8 DUPLICATES REMOVED)

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ANSWER 1 OF 11 SCISEARCH COPYRIGHT 2003 THOMSON ISI on STNDUPLICATE 1

2003:472814 SCISEARCH ACCESSION NUMBER:

THE GENUINE ARTICLE: 683UD

Involvement of AMP-activated protein kinase in glucose TITLE:

uptake stimulated by the globular domain of

adiponectin in primary rat adipocytes

Wu X D; Motoshima H; Mahadev K; Stalker T J; Scalia R; **AUTHOR:** 

Goldstein B J (Reprint)

Thomas Jefferson Univ, Jefferson Med Coll, Div Endocrinol CORPORATE SOURCE:

Diabet & Metab Dis, Dorrance H Hamilton Res Labs, Dept Med, Rm 349 Alumni Hall, 1020 Locust St, Philadelphia, PA 19107 USA (Reprint); Thomas Jefferson Univ, Jefferson Med

Coll, Div Endocrinol Diabet & Metab Dis, Dorrance H Hamilton Res Labs, Dept Med, Philadelphia, PA 19107 USA; Thomas Jefferson Univ, Jefferson Med Coll, Dept Physiol,

Philadelphia, PA 19107 USA

COUNTRY OF AUTHOR: USA

DIABETES, (JUN 2003) Vol. 52, No. 6, pp. 1355-1363. SOURCE:

Publisher: AMER DIABETES ASSOC, 1701 N BEAUREGARD ST,

ALEXANDRIA, VA 22311-1717 USA.

ISSN: 0012-1797. Article; Journal

DOCUMENT TYPE:

English

LANGUAGE:

REFERENCE COUNT:

\*ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS\*

Adiponeetin is an abundant adipocyte-derived plasma protein with AB anti-atherosclerotic and insulin-sensitizing properties that, suppresses hepatic glucose production and enhances glucose uptake into skeletal muscle. To characterize the potential effects of

adiponectin on glucose uptake into adipose cells, we incubated isolated epididyinal rat adipocytes with the globular domain of recombinant adiponectin purified from an E. coli

expression, system. Globular adiponectin increased glucose uptake in adipocytes without stimulating tyrosine phosphorylation of the insulin receptor or insulin receptor substrate-1, and without enhancing phosphorylation of Akt on Ser-473. Globular adiponectin further enhanced, insulin-stimulated glucose uptake at submaximal insulin

concentrations and reversed the inhibitory effect of tumor necrosis, factor-a on insulin-stimulated glucose uptake. Cellular treatment

with globular adiponectin increased the Thr-172 phosphorylation and catalytic activity of AMP-activated protein kinase and enhanced the Ser-79 phosphorylation of acetyl CoA carboxylase; an enzyme downstream of AMP kinase in Adipose cells. Inhibition of AMP kinase activation using two

pharmacological inhibitors (adenine 9-beta-D-axabinofuranoside and compound C) completely abrogated the increase in glucose uptake. stimulated by globular adiponectin, indicating that AMP kinase is integrally involved in the adiponectin signal transduction pathway. Coupled with recent evidence that the effects of adiponectin are mediated via AMP kinase activation in liver and skeletal muscle; the findings reported here provide an important mechanistic link in the signaling effects of adiponectin in

diverse metabolically responsive tissues.

ANSWER 2 OF 11 MEDLINE on STN ACCESSION NUMBER: 2003041047 MEDLINE

PubMed ID: 12547549 DOCUMENT NUMBER: 22436055

Adiponectin and protection against type 2 TITLE:

diabetes mellitus.

COMMENT: Erratum in: Lancet. 2002 Mar 22;361(9362):1060

Spranger Joachim; Kroke Anja; Mohlig Matthias; Bergmann AUTHOR: Manuela M; Ristow Michael; Boeing Heiner; Pfeiffer Andreas н

CORPORATE SOURCE: Department of Nutrition, Endocrinology, and Metabolism,

Benjamin Franklin Medical Centre, Free University Berlin,

Germany.. spranger@mail.dife.de

SOURCE: LANCET, (2003 Jan 18) 361 (9353) 226-8.

Journal code: 2985213R. ISSN: 0140-6736.

PUB. COUNTRY: England: United Kingdom

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Abridged Index Medicus Journals; Priority Journals

ENTRY MONTH: 200302

ENTRY DATE: Entered STN: 20030129

Last Updated on STN: 20030410.

Entered Medline: 20030205

Adiponectin is an adipocyte-derived peptide, which has anti-inflammatory and insulin-sensitising properties. We designed a nested case-control study to assess whether baseline adiponectin concentrations in plasma are independently associated with risk of type 2 diabetes. We found that adiponectin concentrations in plasma were lower among individuals who later developed type 2 diabetes than among controls (mean 5.34 microg/mL [SD 3.49] vs 6.87 microg/mL [4.58], p<0.0001). High concentrations of adiponectin were associated with a substantially reduced relative risk of type 2 diabetes after adjustment for age, sex, waist-to-hip ratio, body-mass index, smoking, exercise, alcohol consumption, education, and glycosylated haemoglobin A(1c) (odds ratio 4th vs 1st quartile 0.3 [95% CI 0.2-0.7], p=0.0051). We conclude that adiponectin is independently associated with a reduced risk of type 2 diabetes in apparently healthy individuals.

L4 ANSWER 3 OF 11 CAPLUS COPYRIGHT 2003 ACS on STN DUPLICATE 2

ACCESSION NUMBER: 2002:646135 CAPLUS

DOCUMENT NUMBER: 137:346864

TITLE: Oligomerization state-dependent activation of

NF-.kappa.B signaling pathway by adipocyte complement-related protein of 30 kDa (Acrp30)

AUTHOR(S); Tsao, Tsu-Shuen; Murrey, Heather E.; Hug, Christopher;

Lee, David H.; Lodish, Harvey F.

CORPORATE SOURCE: Whitehead Institute for Biomedical Research,

whitehead institute for blomedical Resear

Cambridge, MA, 02142, USA

SOURCE: Journal of Biological Chemistry (2002), 277(33),

29359-29362

CODEN: JBCHA3; ISSN: 0021-9258

PUBLISHER: American Society for Biochemistry and Molecular

Biology

DOCUMENT TYPE: Journal LANGUAGE: English

Adipocyte complement-related protein of 30 kDa (Acrp30)/ adiponectin is an adipocyte-derived hormone that affects lipid and glucose metab. in muscle and liver, but its phys. and biochem. properties are poorly characterized. Here we have used several approaches to show that Acrp30 expressed in and purified from Escherichia coli and human embryonic kidney 293T cells forms trimers and hexamers; 293T cells also produce a higher mol. wt. species. Similar Acrp30 oligomers were found in mouse serum as well as in 3T3-L1 adipocyte-conditioned medium, although in different proportions. parallel, we assessed whether Acrp30 is a signaling mol. by searching for promoter or enhancer elements that respond to Acrp30 or its isolated trimeric globular C-terminal domain, gAcrp30. Acrp30 addn. to C2C12 myocytes or myotubes led to activation of NF-.kappa.B transcription factor in a manner dependent upon phosphorylation and degrdn. of I.kappa.B-.alpha.. Importantly, only hexameric and larger isoforms of Acrp30 activated NF-.kappa.B; trimeric Acrp30 or gAcrp30 could not activate NF-.kappa.B. Our data indicate that oligomerization of Acrp30 is important for at least some of its biol. activities, and changes

in the relative abundance of each oligomeric isoform in plasma may

regulate Acrp30 activity.

THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 17

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 4 OF 11 SCISEARCH COPYRIGHT 2003 THOMSON ISI on STN

2002:526350 SCISEARCH ACCESSION NUMBER:

THE GENUINE ARTICLE: 557XP

Purification of globular form of TITLE:

adiponectin in human serum and its potential role

in insulin sensitivity

Waki H (Reprint); Yamauchi T; Kamon J; Ito Y; Uchida S; AUTHOR:

Oike Y; Yamamura K; Kimura S; Kadowaki T

DIABETES, (JUN 2002) Vol. 51, Supp. [2], pp. A455-A455. MA SOURCE:

Publisher: AMER DIABETES ASSOC, 1660 DUKE ST, ALEXANDRIA,

VA 22314 USA. ISSN: 0012-1797. Conference; Journal

DOCUMENT TYPE:

LANGUAGE: English

REFERENCE COUNT:

SCISEARCH COPYRIGHT 2003 THOMSON ISI on STN ANSWER 5 OF 11

2001:647994 SCISEARCH ACCESSION NUMBER:

THE GENUINE ARTICLE: 461BX

Identification and adipocyte differentiation-dependent

expression of the unique disialic acid residue in an

adipose tissue-specific glycoprotein, adipo Q

Sato C; Yasukawa Z; Honda N; Matsuda T; Kitajima K AUTHOR:

(Reprint)

Nagoya Univ, Grad Sch Bioagr Sci, Dept Appl Mol Biosci, CORPORATE SOURCE:

Nagoya, Aichi 4648601, Japan (Reprint); Nagoya Univ, Biosci Ctr, Div Oncogenesis, Dept Anim Sci, Nagoya, Aichi

4648601, Japan

COUNTRY OF AUTHOR:

Japan

JOURNAL OF BIOLOGICAL CHEMISTRY, (3 AUG 2001) Vol. 276, SOURCE:

No. 31, pp. 28849-28856.

Publisher: AMER SOC BIOCHEMISTRY MOLECULAR BIOLOGY INC,

9650 ROCKVILLE PIKE, BETHESDA, MD 20814 USA.

ISSN: 0021-9258.

DOCUMENT TYPE:

Article; Journal English

LANGUAGE: REFERENCE COUNT:

\*ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS\*

Recently, we have shown that alpha2,8-linked disialic acid (diSia) residue occurs in glycoproteins more frequently than ever recognized (Sato, C., Fukuoka, H., Ohta, K., Matsuda, T., Koshino, R., Kobayashi K., Troy, F. A., II, and Kitajima, K. (2000) J. BioL Chem. 275,15422-15431). In the course of identification of the diSia-containing glycoproteins in mammals, the 30-kDa glycoprotein was found in bovine serum. The 30-kDa glycoprotein was shown to be the bovine adipo Q, an adipocyte-specific protein, based on the partial amino acid sequences and the immuno-cross-reactivity with the recombinant mouse adipo Q. The bovine adipo, Q was shown to have no N-linked but O-linked glycan(s) containing the diSia epitope, Neu5Ac alpha2-8Neu5Ac alpha2 --> 3Gal. Furthermore, the diSia epitope was also found in the mouse adipo Q in serum as well as in the 3T3-L1 cells that are fully differentiated into adipocytes. Notably, among the known alpha2,8-sialyltransferases, only the alpha2,8sialyltransferase III mRNA was detected in the 3T3-L1 cells at any stages of differentiation, and the recombinant alpha2,8-sialyltransferase III could sialylate the purified bovine adipo Q. Thus, this study clearly provides the new findings that adipo Q is the diSia-containing qlycoprotein and a physiological substrate of alpha2,8-sialyltransferase III, whose substrates have not been identified so far.

ANSWER 6 OF 11 SCISEARCH COPYRIGHT 2003 THOMSON ISI on STNDUPLICATE 3 L4

2002:21442 SCISEARCH ACCESSION NUMBER:

THE GENUINE ARTICLE: 503WC

Endogenous glucose production is inhibited by the TITLE:

· adipose-derived protein Acrp30

Combs T P; Berg A H; Obici S; Scherer P E; Rossetti L AUTHOR:

(Reprint)

Albert Einstein Coll Med, Dept Pharmacol, 1300 Morris Pk CORPORATE SOURCE:

Ave, Bronx, NY 10461 USA (Reprint); Albert Einstein Coll Med, Dept Cell Biol, Bronx, NY 10461 USA; Albert Einstein Coll Med, Dept Med, Bronx, NY 10461 USA; Albert Einstein Coll Med, Ctr Diabet Res & Training, Bronx, NY 10461 USA; Albert Einstein Coll Med, Dept Mol Pharmacol, Bronx, NY

10461 USA

COUNTRY OF AUTHOR:

USA SOURCE:

JOURNAL OF CLINICAL INVESTIGATION, (DEC 2001) Vol. 108,

No. 12, pp. 1875-1881.

Publisher: AMER SOC CLINICAL INVESTIGATION INC, 35 RESEARCH DR, STE 300, ANN ARBOR, MI 48103 USA.

ISSN: 0021-9738. Article; Journal

DOCUMENT TYPE: LANGUAGE:

English

REFERENCE COUNT: 38

\*ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS\*

Intraperitoneal injection of purified recombinant Acrp30 lowers glucose levels in mice. To gain insight into the mechanism(s) of this hypoglycemic effect, purified recombinant Acrp30 was infused in conscious mice during a pancreatic euglycemic clamp. In the presence of physiological hyperinsulinemia, this treatment increased circulating Acrp30 levels by approximately twofold and stimulated glucose metabolism. The effect of Acrp30 on in vivo insulin action was completely accounted for by a 65% reduction in the rate of glucose production. Similarly, glucose flux through glucose-6-phosphatase (G6Pase) decreased with Acrp30, whereas the activity of the direct pathway of glucose-6-phosphate biosynthesis, an index of hepatic glucose phosphorylation, increased significantly. Acrp30 did not affect the rates of glucose uptake, glycolysis, or glycogen synthesis. These results indicate that an acute increase in circulating Acrp30 levels lowers hepatic glucose production without affecting peripheral glucose uptake. Hepatic expression of the gluconeogenic enzymes phosphoenolpyruvate carboxykinase and G6Pase mRNAs was reduced by more than 50% following Acrp30 infusion compared with vehicle infusion. Thus, a moderate rise in circulating levels of the adipose-derived protein Acrp30 inhibits both the expression of hepatic gluconeogenic enzymes and the rate of endogenous

ANSWER 7 OF 11 SCISEARCH COPYRIGHT 2003 THOMSON ISI on STNDUPLICATE 4

ACCESSION NUMBER: 2001:855020 SCISEARCH

THE GENUINE ARTICLE: 485EK

glucose production.

TITLE: The adipocyte-secreted protein Acrp30 enhances hepatic

insulin action

Berg A H; Combs T P; Du X L; Brownlee M; Scherer P AUTHOR:

(Reprint)

Albert Einstein Coll Med, Dept Cell Biol, Bronx, NY 10467 CORPORATE SOURCE:

> USA (Reprint); Albert Einstein Coll Med, Dept Med, Bronx, NY 10467 USA; Albert Einstein Coll Med, Dept Pathol, Bronx, NY 10467 USA; Albert Einstein Coll Med, Ctr Diabet

Res & Training, Bronx, NY 10467 USA

COUNTRY OF AUTHOR:

SOURCE:

NATURE MEDICINE, (AUG 2001) Vol. 7, No. 8, pp. 947-953. Publisher: NATURE AMERICA INC, 345 PARK AVE SOUTH, NEW

YORK, NY 10010-1707 USA.

ISSN: 1078-8956.

DOCUMENT TYPE: Article; Journal

English LANGUAGE:

REFERENCE COUNT:

\*ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS\*

Acrp30 is a circulating protein synthesized in adipose tissue. A single AB injection in mice of purified recombinant Acrp30 leads to a 2-3-fold elevation in circulating Acrp30 levels, which triggers a transient decrease in basal glucose levels. Similar treatment in ob/ob, NOD (non-obese diabetic) or streptozotocin-treated mice transiently abolishes hyperglycemia. This effect on glucose is not associated with an increase in insulin levels. Moreover, in isolated hepatocytes, Acrp30 increases the ability of sub-physiological levels of insulin to suppress glucose production. We thus propose that Acrp30 is a potent insulin enhancer linking adipose tissue and whole-body glucose metabolism.

ANSWER 8 OF 11 BIOSIS COPYRIGHT 2003 BIOLOGICAL ABSTRACTS INC. on STN

ACCESSION NUMBER: DOCUMENT NUMBER:

2001:278603 BIOSIS PREV200100278603

TITLE:

Adiponectin, an adipocyte-specific secretory protein, inhibits B lymphopoiesis in culture.

AUTHOR (S):

Yokota, Takafumi (1); Oritani, Kenji; Kouro, Taku (1); Meka, Reddy (1); Medina, Kay L. (1); Tomiyama, Yoshiaki;

Matsuzawa, Yuji; Kincade, Paul W. (1)

CORPORATE SOURCE:

(1) Oklahoma Medical Research Foundation, 825 Northeast

13th Street, Oklahoma City, OK, 73104 USA

SOURCE:

FASEB Journal, (March 7, 2001) Vol. 15, No. 4, pp. A318.

print.

Meeting Info.: Annual Meeting of the Federation of American Societies for Experimental Biology on Experimental Biology

2001 Orlando, Florida, USA March 31-April 04, 2001

ISSN: 0892-6638.

DOCUMENT TYPE:

Conference English

LANGUAGE: SUMMARY LANGUAGE: English

The stromal cells that support blood cell production within bone marrow are pre-adipocytes and functional interactions with marrow fat cells have long been suspected. Adiponectin was recently isolated as an adipocyte product and shown to have structural similarities to C1q as well as members of the TNF superfamily. It suppresses myeloid differentiation in short term bone marrow cultures and also inhibits macrophage functions. These observations raised the possibility that precursors of other blood cell lineages interact with fat cells in marrow via adiponectin. We have now determined that the factor blocks B lymphopoiesis in Whitlock-Witte type bone marrow cultures, but not the production of myeloid cells in Dexter cultures. Several observations suggest that non-lymphoid cells represent the target of this new mediator, and the B lymphoid lineage is only indirectly influenced. Highly purified lymphocyte precursors in stromal cell-free, serum-free cultures were unaffected by adiponectin. Similarly, there was no influence on IL-7 responding pro-B cells in clonal assays. The cytokine dramatically inhibited, and even reversed adipogenesis in culture, suggesting that it may normally be a feedback inhibitor of this process. PCR analyses are being conducted with cloned stromal cells that are responsive to adiponectin, with a view to learning if a negative regulator of B lymphopoiesis is induced. Preliminary results suggest that expression of TNFalpha, TGFbeta, interferons and a new interferon-like cytokine known as limitin are not up-regulated by adiponectin. Further studies should be informative about the role of fat cells within bone marrow and could reveal some involvement of adiponectin with lymphocyte production.

ANSWER 9 OF 11 MEDLINE on STN ACCESSION NUMBER: 2000072595 MEDLINE

PubMed ID: 10604883 20072595 DOCUMENT NUMBER:

TITLE: Novel modulator for endothelial adhesion molecules:

adipocyte-derived plasma protein adiponectin.

AUTHOR: Ouchi N; Kihara S; Arita Y; Maeda K; Kuriyama H; Okamoto Y;

Hotta K; Nishida M; Takahashi M; Nakamura T; Yamashita S;

Funahashi T; Matsuzawa Y

CORPORATE SOURCE: Department of Internal Medicine and Molecular Science,

Graduate School of Medicine, Osaka University, Osaka,

Japan.. ouchi@imed2.med.osaka-u.ac.jp

SOURCE: CIRCULATION, (1999 Dec 21-28) 100 (25) 2473-6.

Journal code: 0147763. ISSN: 0009-7322.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Abridged Index Medicus Journals; Priority Journals

ENTRY MONTH: 200001

ENTRY DATE: Entered STN: 20000124

Last Updated on STN: 20000124 Entered Medline: 20000113

BACKGROUND: Among the many adipocyte-derived endocrine factors, we recently found an adipocyte-specific secretory protein, adiponectin, which was decreased in obesity. Although obesity is associated with increased cardiovascular mortality and morbidity, the molecular basis for the link between obesity and vascular disease has not been fully clarified. The present study investigated whether adiponectin could modulate endothelial function and relate to coronary disease. METHODS AND RESULTS: For the in vitro study, human aortic endothelial cells (HAECs) were preincubated for 18 hours with the

indicated amount of adiponectin, then exposed to tumor necrosis factor-alpha (TNF-alpha) (10 U/mL) or vehicle for the times indicated. The adhesion of human monocytic cell line THP-1 cells to HAECs was determined by adhesion assay. The surface expression of vascular cell adhesion molecule-1 (VCAM-1), endothelial-leukocyte adhesion molecule-1 (E-selectin), and intracellular adhesion molecule-1 (ICAM-1) was measured

by cell ELISA. Physiological concentrations of adiponectin dose-dependently inhibited TNF-alpha-induced THP-1 adhesion and expression of VCAM-1, E-selectin, and ICAM-1 on HAECs. For the in vivo study, the concentrations of adiponectin in human plasma were determined by a sandwich ELISA system that we recently developed. Plasma adiponectin concentrations were significantly lower in patients

with coronary artery disease than those in age- and body mass index-adjusted control subjects. CONCLUSIONS: These observations suggest that adiponectin modulates endothelial inflammatory response and that the measurement of plasma adiponectin levels may be helpful

in assessment of CAD risk.

L4 ANSWER 10 OF 11 MEDLINE ON STN ACCESSION NUMBER: 1999194557 MEDLINE

DOCUMENT NUMBER: 99194557 PubMed ID: 10092513

TITLE: Paradoxical decrease of an adipose-specific protein,

adiponectin, in obesity.

AUTHOR: Arita Y; Kihara S; Ouchi N; Takahashi M; Maeda K; Miyagawa J; Hotta K; Shimomura I; Nakamura T; Miyaoka K; Kuriyama H;

J; Hotta K; Shimomura I; Nakamura T; Miyaoka K; Kuriyama H; Nishida M; Yamashita S; Okubo K; Matsubara K; Muraguchi M;

Ohmoto Y; Funahashi T; Matsuzawa Y

CORPORATE SOURCE: Graduate School of Medicine, Institute for Molecular and

Cellular Biology, Osaka University, 2-2 Yamadaoka, Suita,

Osaka, 565-0871, Japan.

SOURCE: BIOCHEMICAL AND BIOPHYSICAL RESEARCH COMMUNICATIONS, (1999

Apr 2) 257 (1) 79-83.

Journal code: 0372516. ISSN: 0006-291X.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199905

ENTRY DATE: Entered STN: 19990525

Last Updated on STN: 20000303

Entered Medline: 19990511

We isolated the human adipose-specific and most abundant gene transcript, apM1 (Maeda, K., et al., Biochem. Biophys. Res. Commun. 221, 286-289, 1996). The apM1 gene product was a kind of soluble matrix protein, which we named adiponectin. To quantitate the plasma adiponectin concentration, we have produced monoclonal and polyclonal antibodies for human adiponectin and developed an enzyme-linked immunosorbent assay (ELISA) system. Adiponectin was abundantly present in the plasma of healthy volunteers in the range from 1.9 to 17.0 mg/ml. Plasma concentrations of adiponectin in obese subjects were significantly lower than those in non-obese subjects, although adiponectin is secreted only from adipose tissue. The ELISA system developed in this study will be useful for elucidating the physiological and pathophysiological role of adiponectin in humans.

Copyright 1999 Academic Press.

L4 ANSWER 11 OF 11 MEDLINE ON STN ACCESSION NUMBER: 97103474 MEDLINE

DOCUMENT NUMBER: 97103474 PubMed ID: 8947845

TITLE: Isolation and characterization of

GBP28, a novel gelatin-binding protein purified

from human plasma.

AUTHOR: Nakano Y; Tobe T; Choi-Miura N H; Mazda T; Tomita M

CORPORATE SOURCE: Department of Physiological Chemistry, School of

Pharmaceutical Sciences, Showa University..

yanakano@pharm.showa-u.ac.jp

SOURCE: JOURNAL OF BIOCHEMISTRY, (1996 Oct) 120 (4) 803-12.

Journal code: 0376600. ISSN: 0021-924X.

PUB. COUNTRY: Japan

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199703

ENTRY DATE: Entered STN: 19970327

Last Updated on STN: 20000303 Entered Medline: 19970317

By use of its affinity to gelatin-Cellulofine, a novel protein, GBP28 AB (gelatin-binding protein of 28 kDa), was obtained from human plasma. GBP28 bound to gelatin-Cellulofine could be eluted with 1 M NaCl. By analysis of its amino-terminal amino acid sequences and the peptides obtained by protease digestion, GBP28 was identified as a novel protein. After repeated gel chromatography of the 1 M NaCl eluate from gelatin-Cellulofine, about 50 micrograms of GBP28 was purified from 500 ml of human plasma. On gel chromatography, the protein migrated as a molecule of about 420 kDa. On SDS-PAGE, its molecular mass was 28 kDa under reducing conditions and 68 kDa under nonreducing conditions. Recently, human mRNA specific to adipose tissue, cDNA clone apM1, has been registered [Maeda, K., Okubo, K., Shimomura, I., Funahashi, T., Matsuzawa, Y., and Matsubara, K. (1996) Biochem. Biophys. Res. Commun. 221, 286-289]. The assumed amino acid sequence of cDNA clone apM1 contained all the sequences of GBP28 and its peptides. Therefore, it is evident that the cDNA clone apM1 encodes GBP28 and the protein is specific to adipose tissue. The clone encodes a polypeptide of 244 amino acids with a secretory signal sequence at the amino terminus, a small non-helical region, a stretch of 22 collagen repeats and a globular domain. Thus, GBP28 appears to belong to a family of proteins possessing a collagen-like domain through which they form homo-trimers, which further combine to make oligomeric complexes. Although its biological function is presently unclear, its adipocyte-specific expression suggests that GBP28 may function as an endogenous factor involved in lipid catabolism and storage

or whole body metabolism.

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END OF SEAL	RCH HISTORY	

## WEST

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## Search Results - Record(s) 1 through 19 of 19 returned.

☐ 1. Document ID: US 20030108883 A1

L1: Entry 1 of 19

File: PGPB

Jun 12, 2003

PGPUB-DOCUMENT-NUMBER: 20030108883

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20030108883 A1

TITLE: Methods for identifying compounds that inhibit or reduce PTP1B expression

PUBLICATION-DATE: June 12, 2003

INVENTOR-INFORMATION:

NAME	CITY	STATE	COUNTRY	RULE-47
Rondinone, Cristina M.	Libertyville	IL	US	
Trevillyan, James M.	Grayslake	IL	US	
Zinker, Bradley A.	Vernon Hills	IL .	US	
Waring, Jeffrey F.	Franklin	MI .	us ·	
Jirousek, Mike	San Diego	CA	US	
Butler, Madeline M.	Santa Fe	CA	US	
Cowsert, Lex M.	Pittsburgh	PA	US	
Monia, Brett P.	Encinitas	CA	US	
Wyatt, Jacqueline	Encinitas	CA	US	

US-CL-CURRENT: 435/6; 435/7.9

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims KWC Draw Desc Image

☐ 2. Document ID: US 20030092736 A1

L1: Entry 2 of 19.

File: PGPB

May 15, 2003

PGPUB-DOCUMENT-NUMBER: 20030092736

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20030092736 A1

TITLE: Substituted azole acid derivatives useful as antidiabetic and antiobesity

agents and method

PUBLICATION-DATE: May 15, 2003

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47 Cheng, Peter T. Princeton · ŊJ US Zhang, Hao Belle Mead NJ US PA Hariharan, Narayanan Richboro US

US-CL-CURRENT: 514/333; 514/340, 514/365, 514/374, 514/396, 546/256, 546/270.4, 546/271.4, 546/272.7, 546/276.4

Full Title Citation Front Review Classification Date Reference Sequences Attachments Claims KNNC Draw Desc Image

3. Document ID: US 20030044396 A1

L1: Entry 3 of 19

File: PGPB

Mar 6, 2003

PGPUB-DOCUMENT-NUMBER: 20030044396

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20030044396 A1

TITLE: Methods for treating diseases and increasing longevity

PUBLICATION-DATE: March 6, 2003

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47

Elia, James P. Scottsdale AZ US

US-CL-CURRENT: 424/93.21; 435/366

Full Title Citation Front Review Classification Date Reference Sequences Attachments KMIC Draw Desc Image

4. Document ID: US 20020132773 A1

L1: Entry 4 of 19 File: PGPB Sep 19, 2002

PGPUB-DOCUMENT-NUMBER: 20020132773

PGPUB-FILING-TYPE: new

DOCUMENT-IDENTIFIER: US 20020132773 A1

TITLE: Methods for reducing fat by administration of adiponectin

PUBLICATION-DATE: September 19, 2002

INVENTOR-INFORMATION:

NAME CITY STATE COUNTRY RULE-47

Kincade, Paul W. Nichols Hill OK US Yokuta, Takafumi Norman OK US

US-CL-CURRENT: 514/12

Full Title Citation Front Review Classification Date Reference Sequences Attachments | KMC | Draw Desc | Image |

5. Document ID: US 6582909 B1

L1: Entry 5 of 19 File: USPT Jun 24, 2003

US-PAT-NO: 6582909

DOCUMENT-IDENTIFIER: US 6582909 B1

TITLE: APM1 biallelic markers and uses thereof

Full Title Citation Front Review Classification Date Reference Sequences Attachments KWIC Draw Desc Image

☐ 6. Document ID: US 6566332 B2 L1: Entry 6 of 19 File:	USPT May 20, 2003
US-PAT-NO: 6566332 DOCUMENT-IDENTIFIER: US 6566332 B2	
TITLE: OBG3 globular head and uses thereof	for decreasing body mass
Full Title Citation Front Review Classification Date Reference	Sequences Attachments KWMC Draw Desc Image
7. Document ID: US 6479238 B1 L1: Entry 7 of 19 File:	USPT Nov 12, 2002
US-PAT-NO: 6479238 DOCUMENT-IDENTIFIER: US 6479238 B1	
TITLE: Polymorphic markers of the LSR gene	
Full Title Citation Front Review Classification Date Reference	Sequences Attachments   KMC   Draw Desc   Image
<pre>\$\bigcup 8. Document ID: US 6344441 B1 L1: Entry 8 of 19 File:</pre>	USPT Feb 5, 2002
US-PAT-NO: 6344441 DOCUMENT-IDENTIFIER: US 6344441 B1 ** See image for Certificate of Correction	**
TITLE: Lipoprotein-regulating medicaments	
Full Title Citation Front Review Classification Date Reference	Sequences Attachments   KMC   Draw Desc   Image
9. Document ID: JP 2002363094 A L1: Entry 9 of 19 File:	JPAB Dec 18, 2002
PUB-NO: JP02002363094A DOCUMENT-IDENTIFIER: JP 2002363094 A TITLE: HEPATIC FIBROGENETIC SUPPRESSOR	
Full Title Citation Front Review Classification Date Reference	Sequences Attachments KWWC Draws Desc Image
☐ 10. Document ID: JP 2000256208 A L1: Entry 10 of 19 File:	JPAB Sep 19, 2000

PUB-NO: JP02000256208A

DOCUMENT-IDENTIFIER: JP 2000256208 A

TITLE: ANTI-INFLAMMATORY AGENT AND PROPAGATION SUPPRESSOR FOR MONOCYTOID CELL

☐ 11. Document ID: WC	3016906 A1	•
L1: Entry 11 of 19	File: EPAB	Feb 27, 2003
	906 A1 OR MONITORING SACCHAROMETABOL	IC ERROR  NAMC Drawn Desc   Image
☐ 12. Document ID: WC	2100427 A1	
L1: Entry 12 of 19	File: EPAB	Dec 19, 2002
JB-NO: WO002100427A1 CCUMENT-IDENTIFIER: WO 2100 ITLE: LIVER GENERATION PROM		
Full   Title   Citation   Front   Review   Class	ification   Date   Reference   Sequences   Attachments	KNAC   Draw Desc   Image
☐ 13. Document ID: WC	2072149 A1	
L1: Entry 13 of 19	File: EPAB	Sep 19, 2002
	149 A1 FAT BY ADMINISTRATION OF ADIP  fication   Data   Reference   Sequences   Attachments	ONECTIN    MMC   Draws Desc   Image
☐ 14. Document ID: WC	) 2061076 A1	
L1: Entry 14 of 19	File: EPAB	Aug 8, 2002
JB-NO: WO002061076A1 CUMENT-IDENTIFIER: WO 2061 TLE: ADIPONECTIN-ASSOCIATE		IQMC   Draw Desc   Image
☐ 15. Document ID: W	2 2022221 (40 + 2	

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TITLE: New primary preadipocyte strain expressing telomerase reverse transcriptase, useful in research applications, screening assays, clinical applications, and in the

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administration of therapeutic agents, particularly for obesity

Full Title Citation Front Review Classification Date Reference Sequences Attachments KWMC Draw Desc Image 16. Document ID: WO 2003016906 A1 L1: Entry 16 of 19 File: DWPI Feb 27, 2003 DERWENT-ACC-NO: 2003-248408 DERWENT-WEEK: 200324 COPYRIGHT 2003 DERWENT INFORMATION LTD TITLE: Assay of adiponectin (GBP28) in biological samples for the diagnosis and monitoring of errors of sugar metabolism Full Title Citation Front Review Classification Date Reference Sequences Attachments KMC Draw Desc Image 17. Document ID: WO 2002100427 A1 JP 2002363094 A L1: Entry 17 of 19 File: DWPI Dec 19, 2002 DERWENT-ACC-NO: 2003-156922 DERWENT-WEEK: 200315 COPYRIGHT 2003 DERWENT INFORMATION LTD TITLE: Liver generation promoter comprises <a href="mailto:adiponectin">adiponectin</a> useful for treating and preventing liver cirrhosis and chronic hepatitis Full Title Citation Front Review Classification Date Reference Sequences Attachments KWIC Draw Desc Clip Img Image 18. Document ID: US 20020132773 A1 WO 200272149 A1 L1: Entry 18 of 19 File: DWPI Sep 19, 2002 DERWENT-ACC-NO: 2003-128066 DERWENT-WEEK: 200312 COPYRIGHT 2003 DERWENT INFORMATION LTD TITLE: Method for decreasing fat in adipocytes or the number of adipocytes comprises administration of adiponectin to adipocytes or tissue containing them Full Title Citation Front Review Classification Date Reference Sequences Attachments KMIC Draw Desc Image 19. Document ID: WO 200261076 A1 L1: Entry 19 of 19 File: DWPI Aug 8, 2002

DERWENT-ACC-NO: 2002-627480

DERWENT-WEEK: 200267

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TITLE: DNA encoding adiponectin-associated protein which inhibits proliferation of vascular smooth muscle cells, applicable in diagnosis and development of preventives or remedies for arteriosclerosis

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PubMed Services Ouchi N, Kihara S, Arita Y, Maeda K, Kuriyama H, Okamoto Y, Hotta K, Nishida M, Takahashi M, Nakamura T, Yamashita S, Funahashi T, Matsuzawa Y.

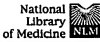
Department of Internal Medicine and Molecular Science, Graduate School of Medicine, Osaka University, Osaka, Japan. ouchi@imed2.med.osaka-u.ac.jp

Related Resources BACKGROUND: Among the many adipocyte-derived endocrine factors, we recently found an adipocyte-specific secretory protein, adiponectin, which was decreased in obesity. Although obesity is associated with increased cardiovascular mortality and morbidity, the molecular basis for the link between obesity and vascular disease has not been fully clarified. The present study investigated whether adiponectin could modulate endothelial function and relate to coronary disease. METHODS AND RESULTS: For the in vitro study, human aortic endothelial cells (HAECs) were preincubated for 18 hours with the indicated amount of adiponectin, then exposed to tumor necrosis factor-alpha (TNF-alpha) (10 U/mL) or vehicle for the times indicated. The adhesion of human monocytic cell line THP-1 cells to HAECs was determined by adhesion assay. The surface expression of vascular cell adhesion molecule-1 (VCAM-1), endothelial-leukocyte adhesion molecule-1 (E-selectin), and intracellular adhesion molecule-1 (ICAM-1) was measured by cell ELISA. Physiological concentrations of adiponectin dose-dependently inhibited TNF-alpha-induced THP-1 adhesion and expression of VCAM-1, E-selectin, and ICAM-1 on HAECs. For the in vivo study, the concentrations of adiponectin in human plasma were determined by a sandwich ELISA system that we recently developed. Plasma adiponectin concentrations were significantly lower in patients with coronary artery disease than those in age- and body mass index-adjusted control subjects. CONCLUSIONS: These observations suggest that adiponectin modulates endothelial inflammatory response and that the measurement of plasma adiponectin levels may be helpful in assessment of CAD risk.

PMID: 10604883 [PubMed - indexed for MEDLINE]







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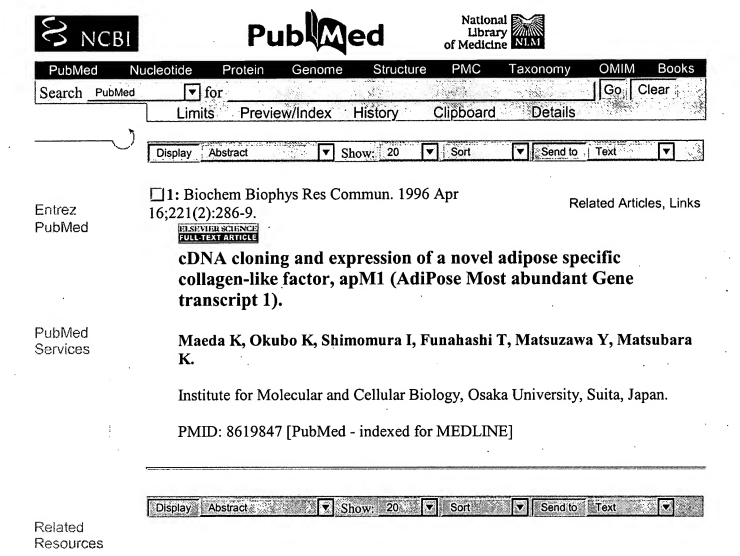
Among them, plasminogen activator-1 (PAI-1), which is a regulator of the fibrinolytic system, was overexpressed in the visceral fat in an animal model of obesity. Plasma levels of PAI-1 were closely correlated with visceral fat adiposity. Thus, PAI-1 secreted from visceral fat may play some role in thrombotic vascular disease in visceral obesity. Adiponectin, a novel adipose-specific gene product, which has a matrix-like structure, is abundantly present in the bloodstream. Dysregulated secretion of adiponectin may be related to vascular disease in obesity. Biologically active molecules secreted from adipose tissue (adipocytokines) may have important roles in the development of atherosclerotic disease in obesity.

## Publication Types:

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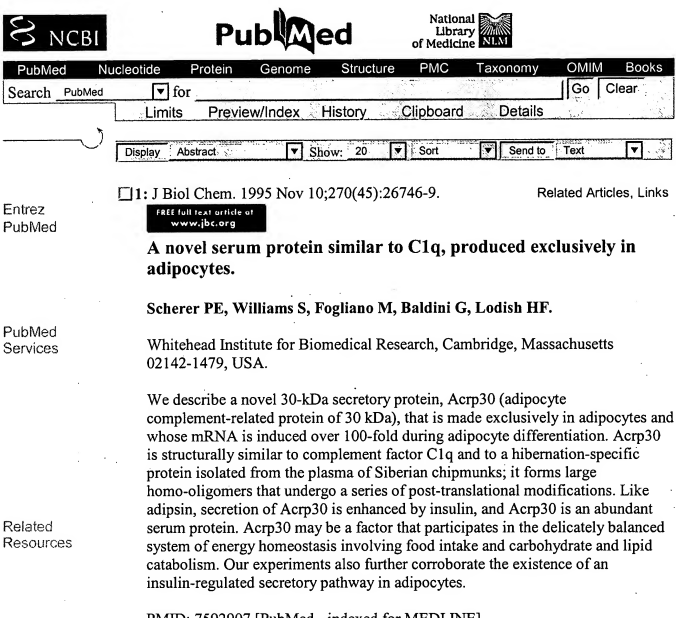






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Entrez PubMed	1: Biochem Biophys Res Commun. 1999 Apr 2;257(1):79-83. Related Articles, Links BUSIEVIER SCIENCE PULLSTEXT ARTICLE  Paradoxical decrease of an adipose-specific protein, adiponectin, in obesity.
PubMed Services	Arita Y, Kihara S, Ouchi N, Takahashi M, Maeda K, Miyagawa J, Hotta K, Shimomura I, Nakamura T, Miyaoka K, Kuriyama H, Nishida M, Yamashita S, Okubo K, Matsubara K, Muraguchi M, Ohmoto Y, Funahashi T, Matsuzawa Y.
	Graduate School of Medicine, Institute for Molecular and Cellular Biology, Osaka University, 2-2 Yamadaoka, Suita, Osaka, 565-0871, Japan.
Related Resources	We isolated the human adipose-specific and most abundant gene transcript, apM1 (Maeda, K., et al., Biochem. Biophys. Res. Commun. 221, 286-289, 1996). The apM1 gene product was a kind of soluble matrix protein, which we named adiponectin. To quantitate the plasma adiponectin concentration, we have produced monoclonal and polyclonal antibodies for human adiponectin and developed an enzyme-linked immunosorbent assay (ELISA) system. Adiponectin was abundantly present in the plasma of healthy volunteers in the range from 1.9 to 17.0 mg/ml. Plasma concentrations of adiponectin in obese subjects were significantly lower than those in non-obese subjects, although adiponectin is secreted only from adipose tissue. The ELISA system developed in this study will be useful for elucidating the physiological and pathophysiological role of adiponectin in humans. Copyright 1999 Academic Press.  PMID: 10092513 [PubMed - indexed for MEDLINE]
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